§314.96

Generic Drugs, 7500 Standish Pl., Rockville, MD 20855.

- (3) This paragraph does not apply to a use patent that claims no uses for which the applicant is seeking approval.
- (b) Sending the notice. The applicant shall send the notice required by paragraph (a) of this section when it receives from FDA an acknowledgment letter stating that its abbreviated new drug application is sufficiently complete to permit a substantive review. At the same time, the applicant shall amend its abbreviated new drug application to include a statement certifying that the notice has been provided to each person identified under paragraph (a) of this section and that the notice met the content requirements under paragraph (c) of this section.
- (c) Contents of a notice. In the notice, the applicant shall cite section 505(j)(2)(B)(ii) of the act and shall include, but not be limited to, the following information:
- (1) A statement that FDA has received an abbreviated new drug application submitted by the applicant containing any required bioavailability or bioequivalence data or information.
- (2) The abbreviated application number
- (3) The established name, if any, as defined in section 502(e)(3) of the act, of the proposed drug product.
- (4) The active ingredient, strength, and dosage form of the proposed drug product.
- (5) The patent number and expiration date, as submitted to the agency or as known to the applicant, of each patent alleged to be invalid, unenforceable, or not infringed.
- (6) A detailed statement of the factual and legal basis of the applicant's opinion that the patent is not valid, unenforceable, or will not be infringed. The applicant shall include in the detailed statement:
- (i) For each claim of a patent alleged not to be infringed, a full and detailed explanation of why the claim is not infringed.
- (ii) For each claim of a patent alleged to be invalid or unenforceable, a full and detailed explanation of the grounds supporting the allegation.

- (7) If the applicant does not reside or have a place of business in the United States, the name and address of an agent in the United States authorized to accept service of process for the applicant.
- (d) Amendment to an abbreviated application. If an abbreviated application is amended to include the certification described in §314.94(a)(12)(i)(A)(4), the applicant shall send the notice required by paragraph (a) of this section at the same time that the amendment to the abbreviated application is submitted to FDA.
- (e) Documentation of receipt of notice. The applicant shall amend its abbreviated application to document receipt of the notice required under paragraph (a) of this section by each person provided the notice. The applicant shall include a copy of the return receipt or other similar evidence of the date the notification was received. FDA will accept as adequate documentation of the date of receipt a return receipt or a letter acknowledging receipt by the person provided the notice. An applicant may rely on another form of documentation only if FDA has agreed to such documentation in advance. A copy of the notice itself need not be submitted to the agency.
- (f) Approval. If the requirements of this section are met, FDA will presume the notice to be complete and sufficient, and it will count the day following the date of receipt of the notice by the patent owner or its representative and by the approved application holder as the first day of the 45-day period provided for in section 505(j)(4)(B)(iii) of the act. FDA may, if the applicant provides a written statement to FDA that a later date should be used, count from such later date.

[59 FR 50366, Oct. 3, 1994, as amended at 68 FR 36705, June 18, 2003; 69 FR 11310, Mar. 10, 2004; 74 FR 9766, Mar. 6, 2009; 74 FR 36605, July 24, 2009]

§ 314.96 Amendments to an unapproved abbreviated application.

(a) Abbreviated new drug application.
(1) An applicant may amend an abbreviated new drug application that is submitted under §314.94, but not yet approved, to revise existing information or provide additional information.

Food and Drug Administration, HHS

Amendments containing bioequivalence studies must contain reports of all bioequivalence studies conducted by the applicant on the same drug product formulation, unless the information has previously been submitted to FDA in the abbreviated new drug application. A complete study report must be submitted for any bioequivalence study upon which the applicant relies for approval. For all other bioequivalence studies conducted on the same drug product formulation as defined in §320.1(g) of this chapter, the applicant must submit either a complete or summary report. If a summary report of a bioequivalence study is submitted and FDA determines that there may be bioequivalence issues or concerns with the product, FDA may require that the applicant submit a complete report of the bioequivalence study to FDA.

- (2) Submission of an amendment containing significant data or information before the end of the initial review cycle constitutes an agreement between FDA and the applicant to extend the initial review cycle only for the time necessary to review the significant data or information and for no more than 180 days.
- (b) The applicant shall submit a field copy of each amendment to §314.94(a)(9). The applicant, other than a foreign applicant, shall include in its submission of each such amendment to FDA a statement certifying that a field copy of the amendment has been sent to the applicant's home FDA district office.

[57 FR 17983, Apr. 28, 1992, as amended at 58 FR 47352, Sept. 8, 1993; 64 FR 401, Jan. 5, 1999; 73 FR 39609, July 10, 2008; 74 FR 2861, Jan. 16, 2009]

§ 314.97 Supplements and other changes to an approved abbreviated application.

The applicant shall comply with the requirements of §§314.70 and 314.71 regarding the submission of supplemental applications and other changes to an approved abbreviated application.

§314.98 Postmarketing reports.

(a) Except as provided in paragraph (b) of this section, each applicant having an approved abbreviated new drug

application under §314.94 that is effective shall comply with the requirements of §314.80 regarding the reporting and recordkeeping of adverse drug experiences.

- (b) Each applicant shall submit one copy of each report required under §314.80 to the Central Document Room, Center for Drug Evaluation and Research, Food and Drug Administration, 5901-B Ammendale Rd., Beltsville, MD 20705-1266.
- (c) Each applicant shall make the reports required under §314.81 and section 505(k) of the act for each of its approved abbreviated applications.

[57 FR 17983, Apr. 28, 1992, as amended at 64 FR 401, Jan. 5, 1999; 74 FR 13113, Mar. 26, 2009]

§ 314.99 Other responsibilities of an applicant of an abbreviated application.

- (a) An applicant shall comply with the requirements of §314.65 regarding withdrawal by the applicant of an unapproved abbreviated application and §314.72 regarding a change in ownership of an abbreviated application.
- (b) An applicant may ask FDA to waive under this section any requirement that applies to the applicant under §§314.92 through 314.99. The applicant shall comply with the requirements for a waiver under §314.90.

Subpart D—FDA Action on Applications and Abbreviated Applications

SOURCE: 50 FR 7493, Feb. 22, 1985, unless otherwise noted. Redesignated at 57 FR 17983, Apr. 28, 1992.

§ 314.100 Timeframes for reviewing applications and abbreviated applications.

- (a) Except as provided in paragraph (c) of this section, within 180 days of receipt of an application for a new drug under section 505(b) of the act or an abbreviated application for a new drug under section 505(j) of the act, FDA will review it and send the applicant either an approval letter under \$314.105 or a complete response letter under \$314.110. This 180-day period is called the "initial review cycle."
- (b) At any time before approval, an applicant may withdraw an application